

who used combination therapy (topical corticosteroid and TCI) (68%). Type of health plan, Medicaid eligibility status, number of therapeutic class, comorbidity, hospitalization or not and AD related costs during 12 months before AD medication started were significantly associated with AD medication adherence. Adherence to AD medication was significantly negatively associated with total annual healthcare costs ( $p < 0.001$ ) and with AD related costs ( $p < 0.001$ ), adjusted for patient demographic, comorbidity, and healthcare utilization characteristics before AD medication started. **CONCLUSIONS:** Poor adherence to topical medication was observed in pediatric AD, and adherence rates differed by the type and combination of AD medication therapy. The detrimental effect of poor adherence on healthcare economic outcome was significant, which implies a need to improve adherence in order to reduce the financial impact of non-adherence. Factors which could contribute non-adherence and financial burden need to be refined and targeted by intervention to improve humanistic and economic outcomes of treatment.

## PSS26

### PATIENT'S EVALUATION OF THE QUICKNESS OF ACTION OF GINGIVAL INFLAMMATION TREATMENTS

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**OBJECTIVES:** Gingivitis is defined as lesions on the gingival margin, expressed through gum redness, bleeding, localized edema, and gingival sensitivity. It is most often caused by substances produced by bacterial plaque, or dental biofilm, which develops along the gingival crevice. To evaluate, using patient's interview results, the quickness of action of several treatments for gingival inflammation. **METHODS:** Observational, prospective, longitudinal, multicentric study carried out in France, using data collected by participating dentists and dental surgeons. **RESULTS:** A total of 316 patients with gingivitis returned their questionnaire. women: 65.25%, current smokers: 22.93% and 28.51% were ex-smokers. Light and heavy bleeding during brushing was reported by 45.74% and 33.33% of patients respectively. 45.70% reported visible redness, 56.34% reported swollen gums, 13.19% had lesions, but above all 50.52% reported pain. Finally, 62.26% had previous history of gingivitis symptoms. As for dental surgeons: 96.36% had performed scaling, 15.15% gingival curettage, and 13.64% radicular resurfacing. A total of 78.4% judged gingival inflammation to be moderate to severe, 63.10% said it had spread (>30%). In terms of treatment: 98.62% gave patients oral hygiene advice, 87.98% advised on brushing methods, 69.91% recommended specific toothpaste, and 78.85% a mouthwash. A total of 30.61% had generalized inflammation after 1 month, reducing to just 11.24% at 2 months and 15.63% at 3 months. A total of 88.08% reported improvement in inflammation after the first month, 91.59% at 2 months and 93.59% at 3 months. A total of 83.93% felt less pain after 1 month of treatment, 87.90% after 2 months and 92.08% after 3 months. ( $p = 0.0418$ ). 89.13% felt their treatment was effective after 1 month, 97.79% after 2 months and 96.15% after 3 months. ( $p = 0.0036$ ). **CONCLUSIONS:** In terms of satisfaction, 86.52% were satisfied after 1 month, 94.85% after 2 months and 95.92% after 3 months. ( $p = 0.0076$ ). 87.57% felt their treatment was easy to follow after 1 month, 86.76% after 2 months and 92.08% after 3 months. Above all, after the first month of treatment, 88.83% said they would continue using the treatment in prevention even after complete disappearance of gingivitis.

## Sensory Systems Disorders – Research on Methods

## PSS27

### EFFECT OF TREATMENT SWITCH ON THE COST-EFFECTIVENESS OF BIOLOGICS IN PSORIASIS IN PERU AND COLOMBIA

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**OBJECTIVES:** To evaluate the effect of treatment switch on the cost-effectiveness of biologics used in patients with moderate or severe psoriasis in Colombia and Peru. **METHODS:** In a previous study (Alandete, JC accepted in the ISPOR 13<sup>th</sup> Annual European Congress) cost effectiveness of etanercept, adalimumab, ustekinumab and infliximab was estimated based on label information for first-(induction) year and second(maintenance) year assuming a 100% treatment continuation (\$1USD=COL\$1.832=SOL\$2.75). For etanercept two induction schemes were considered: 50mg weekly 52 weeks-D1- and 100mg 12 weeks followed by 50 mg 40 weeks-D2-. Effectiveness was evaluated as 75% reduction in Psoriasis Area and Severity Index-PASI 75- infliximab=80%; ustekinumab=69%; adalimumab=59%; etanerceptD2=52%; etanerceptD1=39%. Infliximab and ustekinumab effectiveness were not significantly different. Both were significantly superior to etanercept (Hawkins et al. meta-analysis presented in the 14<sup>th</sup> International ISPOR). In this abstract we developed a new model estimating switching probabilities due to treatment failure at week 12 and adverse events. Biologics costs were adjusted considering time on the pre and post-switching periods. Treatment effectiveness was adjusted when biologics were used after switching due to treatment failure. **RESULTS:** Introduction of switching effect ratified ustekinumab dominance in Colombia (\$US44,675 in 2 years) generating cost savings of -\$US4.049 versus etanerceptD1; -\$US4.049 versus adalimumab; -\$US7.844 versus etanerceptD2 and -\$US27.517 versus infliximab; with higher or same effectiveness than the other biologics in that country. In Peru, ustekinumab changed from being the most cost-effective option and became the dominant option (\$US41.827 in 2 years) generating cost savings of -\$US283 versus etanerceptD1; -\$US489 versus adalimumab; -\$US3581 versus etanerceptD2 and -\$US13.499 versus infliximab. **CONCLUSIONS:** In the studied countries inclusion of the switching effect due to treatment failure and adverse events ratifies cost-savings observed in Colombia and makes ustekinumab the cost-saving option in Peru. These results corroborate those observed in the USA and Europe.

## PSS28

### EFFECT OF DIFFERENT RECALL PERIODS ON DRY EYE SYMPTOM RATINGS

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**OBJECTIVES:** Clinical studies of dry eye disease (DED), a highly symptomatic disease, often ask patients to evaluate their DED symptoms using patient-reported outcomes instruments. Most of these instruments use a one-week recall period. The effect of this recall period on the accuracy of DED symptom assessments has not been documented. The purpose of our research was to compare self-reported DED symptoms between one-week and daily recall periods. **METHODS:** We enrolled 156 DED patients to a web-based observational study to assess their DED symptoms once a day for 9 days. For each of the 14 symptoms, we asked the patients to rate the frequency and intensity on a 0-6 rating scale, with a higher score indicating worse symptom. The assessments on Days 1 and 9 had a one-week recall period, while the assessments on Days 2-8 had a one-day recall period. We then calculated the mean weekly scores for Day 1 and Day 9 and the mean daily scores for Days 2-8, and tested the differences between the mean weekly and daily scores using matched-pair t tests without multiplicity adjustment. **RESULTS:** The Day 1 mean weekly scores were significantly higher than the mean daily scores for all 14 symptoms in both frequency and intensity. The Day 1 mean weekly scores were also significantly higher than the Day 9 mean weekly scores in 10 frequency and 11 intensity scores. The Day 9 mean weekly scores were slightly higher than the daily scores; however, most of the differences were not statistically significant. **CONCLUSIONS:** Patients' self-ratings of their DED symptoms using a one-week recall period are consistently inflated when compared to their ratings using a one-day recall period. Such inflation should be considered when designing clinical studies for DED.

## PSS29

### DEVELOPMENT OF THE MODIFIED OCULAR COMFORT INDEX (MOCI)

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**OBJECTIVES:** Dry eye disease (DED) is characterized by symptoms of ocular discomfort, visual disturbance and reduced tolerance to environmental stressors. DED has a significant negative impact on the quality of life (QOL) of persons affected, and imposes a massive burden on medical resources owing to its high prevalence and chronic nature. It is not known if available patient-reported outcome (PRO) instruments fully capture the scope of DED symptoms and their impact on QOL. The purpose of our ongoing research is to develop a PRO instrument that meets the needs of clinical studies investigating potential treatments for DED. **METHODS:** Patients with DED in five countries (United States, United Kingdom, Spain, Japan and Korea) were interviewed to identify their symptoms and the impact of the disease on QOL (n=120). Based on these results, items were drafted that were tested in two web-based studies with mild-moderate DED subjects (n=106 and 156) and face-to-face interviews with severe DED subjects (n=22). **RESULTS:** Items enquiring about 8 additional symptom experiences (16 items grouped in doublets asking about frequency/intensity) were added to the original Ocular Comfort Index (OCI) using the same question format and response structure (fluctuating vision, light sensitivity, redness, foreign body sensation, excessive tearing, excessive blinking, ocular irritation and stickiness). Additionally, 2 items that enquired about the most bothersome symptom and the extent of bother, and 12 items that appraised how symptoms interfered with the ability to perform daily activities were included. **CONCLUSIONS:** Patient interviews suggest that available PRO instruments do not fully capture the scope of DED symptoms and their impact on QOL. The modified OCI (mOCI) will be used in clinical studies to facilitate its refinement and validation.

## PSS30

### BURDEN OF INFANTILE HEMANGIOMA: DEVELOPMENT OF A QUESTIONNAIRE

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**OBJECTIVES:** Infantile hemangioma (IH) develops during the first weeks of life; it normally forms within 3 to 6 months, then regresses very slowly over a duration of 3 to 7 years. In complicated forms, it is possible to encounter haemorrhaging, necroses and ulcerations, infections and, more exceptionally, respiratory distress, cardiovascular shunt. To explore the handicap, in its largest sense, generated by IH using a questionnaire to express the burden on the daily life of the parents. **METHODS:** The questionnaire was developed following a strict methodological process, involving a multidisciplinary team incorporating various players (doctors, nurses, social workers) who are involved in the treatment of patients or who are specialised in the construction of questionnaires. A review of the literature and discussions with the families were conducted in order to identify the concepts related to the pathology. **RESULTS:** Exploratory assessments showed that the concept of burden could be structured around two main modules: assess the impact directly for the first-module. The consequences of IH on daily life, family and personal relationships, work, financial situation and psychological impact for the second-module. A third module focuses on the behaviour of the child; this module will evolve over time and depending on the analyses. Fifty-six preliminary items were identified following a first discussion. A first analysis managed to reduce these items to 36 whilst conserving the 3 modules but making it easier to use the analysis. **CONCLUSIONS:** The Hemangioma-Burden-Questionnaire will allow clinicians to better understand the impact and consequences of the pathology on the family. It will also allow the development of the burden to be monitored according to the rate of development of the illness and its treatment. It will also allow the